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(54) Activated carbon products and their manufacture

(57) An activated carbon product such as a charcoal cloth or felt has, in addition to any activating material, a metal eg Zn, Al, Ca, Mg or Fe uniformly dispersed therein. The metal may be catalytic or bactericidal, and a particularly preferred product of the invention contains silver and is suitable for use as a surgical dressing.

SPECIFICATION

Activat d carbon produ ts and th ir manufacture

This invention relates to activated carbon products, in particular activated charcoal cloths and felts, and to methods for their manufacture. The products of this invention can be of, for example, bactericidal utility.

Various method for producing activated charcoal cloths are known. For example, GB-A-1455531 discloses impregnating cellulose fibres with a reactive phosphorus compound and heating the impregnated fibres under certain conditions. The Kirk-Othmer Encyclopaedia of Chemical Technology, 16, 3rd Ed., 136 discloses the preparation of novoloid-based activated carbon in a one-step process, combining both carbonisation and activation, in an oxygen-free atmosphere containing steam and/or CO₂, at about 900°C. The products are said to have uniform pore size.

GB-A-1301101 discloses a particularly
useful, and commercially used, process for
preparing activated carbon products in fibrous
form. Rayon, for example, is impregnated
with a solution of inorganic halides, e.g. a
mixture of ammonium, zinc and hexahydrated
aluminium chlorides. The impregnation is followed by a controlled heating stage.

The utility of a carbonised fabric in surgical dressings has been appreciated for over 50 years. GB-A-386067 discloses surgical dressings comprising woven or entangled carbonised fibres. Such dressings are also disclosed as supports for therapeutic or antiseptic materails and it is stated that "the dressings will hold in considerable quantities iodine,

40 formol, lime, oxygen, bacillary toxins, and the like". The use of, say, iodine in the dressings disclosed in GB-A-386067 appears to be a consequence of the adsorptive characteristics of charcoal cloths. Charcoal cloth is an excel-

45 lent adsorbent for organic water-contaminants such as phenol, organic acids and insecticides. Charcoal cloth can also be used, e.g. in gas masks, to remove undesirable gases from the air.

50 EP-A-0053936 discloses surgical dressings comprising activated charcoal, preferably as activated charcoal cloth, impregnated with an anti-microbial agent, iodine being preferred. A characteristic of this disclosure is

55 that no more than 20%, and preferably about 5%, of the adsorptive sites of the activated charcoal are saturated with the anti-microbial ag nt. Such a product probably contains less, say, i dine than an impregnated product as

60 disclosed in GB-A-386067, but still suffers from the disadvantage that i dine is easily r moved from the cloth in the pes nce of aqueous media. It is generally consider d undesirabl that fr iodin should be allowed to

65 c m into contact with a wound, and y t this

can b a probl m ass ciated with the use f surgical dr ssings, containing i dine, as disclosed in EP-A-0053936.

Charcoal cloth has considerable utility of its 70 own as a wound dressing. It can adsorb unpleasant odours of the type which often emanate from infected wounds; in addition, it can adsorb bacteria.

Charcoal cloth may contain traces of ele-75 ments used in the activation procedure. As can be seen, the nature of the product has made it easy to introduce other materials, such as bactericides, subsequently. Charcoal cloth post-impregnated with silver is also

80 known, as a chemisorbent. It would nevertheless be desirable to extend the utility of charcoal cloth, e.g. in medical practice, to take account of its inherent characteristics and to supplement them with properties which are

85 not disadvantaged in the manner described above or in general, following predominantly surface application, by post-impregnation, of a desired additive.

According to the present invention, an acti-90 vated charcoal product has, in addition to any activating material, a metal uniformly dispersed therein.

Metal elements of the compounds used to activate carbon in GB-A-1301101 are Zn,

95 Al, Ca, Mg, Fe (which all have halides with the common, apparently essential, Lewis acid characteristics), Pb, Co and Ba. The metals used in the present invention are intended to provide the product with additional, beneficial

100 properties, e.g. catalytic or bactericidal. Suitable metals for use in the invention are those of the Group VIII elements such as Fe or those of at. no. 76 to 78, e.g. Ir and Pt, and Group Ib, e.g. Ag of Cu.

105 It would obviously save time, labour and cost if, say, charcoal cloth could be impregnated with, say, the bactericidal metal silver, by use of a suitable silver compound at the same time as the activating compounds. For

110 example, a procedural step is saved if the desired metal could be introduced with the halide solution used, before the heating steps, in the procedure of GB-A-1301101, rather than by impregnation after the activated ma-115 terial has been obtained.

However, if a solution of soluble silver salt such as silver nitrate is added to a solution of halides as described in GB-A-1301101, insoluble silver halide is precipitated out. The

soluble silver halide is precipitated out. The 120 result is poor, non-uniform impregnation of the charcoal cloth, or even no impregnation whatsoever, and the presence of potentially undesirable nitrate. A conventienal attempt to overcome this problem, .g. by the addition of

125 ammonia which dissolves and prevents precipitation of silver chloride by forming complexed in second i

130 According to a second aspect of the present

inv nti n, a pr cess f r the pr parati n of an activat d carb n pr duct compris s treating a fibr us carb hydrat mat rial with a s luti n f on or mor L wis acid halid s of Zn, Al,

5 Ca, Mg and Fe; a compound of a further metal element whose halide is relatively insoluble with respect to the, or the mixture of, Lewis acid halides; ammonia; and a sequestering agent; and then drying, carbonising and activating the carbohydrate material. The drying, carbonising and activating may be conducted in conventional manner, e.g. by the procedures described in GB-A-1301101, the contents of which are incorporated herein by 15 reference.

The process of the invention allows the preparation of a product of the invention. The "further" metal element can be any of those described above as having desirable proper20 ties, supplementing those of the activated product, and which have substantially insoluble halides. Added metal element can be uniformly dispersed in the product to give the desired, e.g. catalytic or bactericidal activity.
25 This can be achieved without the precipitation or other problems described above.

The sequestering agent is preferably a hydroxy-carboxylic acid. A suitable hydroxycarboxylic acid is tartaric acid. Citric acid is 30 presently most preferred.

The impregnating solution is suitably prepared by dissolving the Lewis acid or acids, e.g. a mixture of zinc and aluminium chlorides, in water, and then adding the sequest-35 ering agent, ammonia and a soluble salt of the desired metal. An ammoniacal silver halide solution could be used to supply both silver and ammonia.

The quantities of the materials contained in 40 the solution can be determined fairly readily by simple experiment. However as a guide, if the solution contains, by weight, 3% ammonium chloride, 3% zinc chloride and 3% aluminium chloride hexahydrate, as mentioned

45 above, it has been found that the addition of between 3% and 5% by weight citric acid prevents the formation of metal hydroxides when ammonium hydroxide is added to the solution, though greater amounts may be

50 used if desired, particularly if the chloride concentrations are increased. The 3% values for chloride concentration are in fact optimum figures; as little as 2% could be usd. 5% chloride might require about 7–8% citric acid.

Moreover, the amount of ammonium hydroxide required to suppress the formation of silver chloride (when the silver salt is added) is dependent in the amount of silving desired in the final impregnation solution, and can be determined by experiment. If cliudiness is observed in the solution, further ammonia can

While the preferred halides are the chlorides, fluorid s, bromides and iodides can be 65 us d in some cases (though evilution of HF

be added to eliminate precipitation.

during carb nisation is obviously disadvantage us). Agl and AgBr have sufficient subility in ammonia tegive solutions of the required concontration for impregnation (e.g. less than 70 0.1% by weight Ag).

The above description can be generalised when it is desired to disperse a metal other than silver in the product. In determining the amount of the metal which is desired, in

- 75 accordance with the preceding description, the yield of charcoal cloth given by any normal method of manufacture will be generally known or can be easily established. The, say, catalytic or bactericidal effect which is desired
- 80 in the product can be achieved at low levels. Thus, for example, the product will usually comprise at least 0.05 or 0.1, and often at least 0.2, but need not contain more than 5, 2 or even 1, and often no more than 0.5, %
- 85 by weight Ag or other desired metal. A product of the invention can be seen, by suitable microscopic examination, to have a uniform distribution of very small particles of, say silver or silver oxide, extending through the
- 90 thickness of the product. It can be seen quite clearly as distinct from a post-impregnated product, where relatively large agglomerations of the, say, silver or silver oxide are present, and at the surface of the product.
- 95 Other than the presence of the added material, a product of the invention can have all the characteristics associted with activated carbon products. It may be produced in the usual way, e.g. from rayon. It may be a felt or
- 100 a knitted or, typically, woven cloth. A cloth, typically from 0.2 to 1 mm thick, containing uniformly distributed silver, can be advantageously used as a surgical dressing or chemisorbent.
- 105 The method and products of the invention, and their utility, will now be illustrated.

Example 1

To approximately 5 litres of tap water were 110 added:

	ammonium chloride	225 g
	zinc chloride	225 g
	aluminium chloride	225 g
115	citric acid	300 g
	880 ammonia (fresh bottle)	900 cm ³

An aqueous solution containing 15 g silver nitrate in c. 400 cm³ distilled water was made 120 up and acidified with nitric acid (20%; 5 cm³). (This addition is intended to prevent seed crystals of silver chloride forming). This soluti n was k pt stoppered and in a dark place.

125 The silv r nitrat soluti n was added to the bulk liquid (stirred) in aliquots f about 25–50 cm³. Whit pr cipitation quickly disappeared. The addition of the final aliquit produced a persist nt whit pricipitat and a 130 furth r addition f 880 ammonia (200 cm³)

was made. The solution became clear again. Volume was adjust dito 7.51, to give an impregnation solution containing:

5	ammonium chloride	3%
	zinc chloride	3%
	aluminium chloride	3%
	citric acid	4%
	ammonia (as NH ₃)	5%
10	silver nitrate	0.2%

Three lengths of rayon cloth (25 cm × 5 m) were dipped separately into a shallow trough containing the impregnation solution. Dipping 15 time was approximately 2 seconds; rolls following on and being allowed to drain with intermittent turning. Each length was passed through roller nips at 345 kPa and plant oven-dried (at 125°C) by a single pass.

20 A silver analysis was made on a 3 g sample. The analysis was conducted by ashing a sample at 750°C, moistening the resultant ash with concentrated nitric acid, and reigniting the moistened sample to constant weight, in order to ensure that all halides were expelled. The final residue was boiled with 10 cm³ 8 M nitric acid and the entire solution diluted and titrated directly against standardised potassium thiocyanate solution 30 using the Volhard procedure. The silver content was 0.23% by weight.

Example 2

Four pieces of rayon felt (25 × 46 cm) were separately dipped into a shallow trough containing the impregnation solution used in Example 1. Dipping time was approximately 5 seconds. The pieces were rolled consecutively onto a 38 mm diameter tube and drained with intermittent turning. The pieces were oven-dried without pressure by residence of some ten minutes.

All samples were stored in polypropylene sheet and again in black pvc to reduce ultra45 violet penetration pending charring of the samples. All samples were charred at 360°C in carbon dioxide, followed by activation in carbon dioxide at 950°C.

The silver content, by the analysis described 50 in Example 1, was 0.40% by weight.

The cloth and felt products of Examples 1 and 2 have been demonstrated as active against Staphylococcus aureus (Oxford), Bacillus subtilis NCTC 8236, E.coli aeruginosa 55 799 and its envelope mutant 799/61. These

5 799 and its envelope mutant 799/61. These results have been obtained in broth and on agar at dilutions of up to 1:100 (at least for the cloth of Example 1).

Products such as in the Examples can b 60 used in catalysis, .g. the breakdown of arsine and phosphin .

The term "activating" is used herein to describe the se elements, solutions and procedures which are conventionally used in the 65 activation of carbon, although it may only be

a heat-treatment which is conventionally und rst od to pr vid tru activati n. Thus, zinc has been d scrib d as an "activating" element alth ugh its primary r I is to provide 70 strength and flexibility.

CLAIMS

- An activated carbon product having, in addition to any activating material, a metal
 uniformly dispersed therein.
 - 2. A product according to claim 1, in the form of an activated charcoal cloth or felt.
- 3. A product according to claim 1 or claim 2, which comprises from 0.1 to 2% by 80 weight of the metal.
 - 4. A product according to any preceding claim, in which the metal is silver.
- An activated charcoal cloth or felt which comprises from 0.1 to 1% by weight of 85 silver uniformly dispersed therein.
 - A process for the preparation of an activated carbon product, which comprises treating a fibrous carbohydrate material with a solution of one or more Lewis acid halides of
- 90 Zn, Al, Ca, Mg and Fe; a compound of a further metal element whose halide is relatively insoluble with respect to the, or the mixture of, Lewis acid halides; ammonia; and a sequestering agent; and then drying, car-
- 95 bonising and activating the carbohydrate material.
 - 7. A process according to claim 6, in which the sequestering agent is a hydroxycar-boxylic acid.
- 100 8. A process according to claim 7, in which the hydroxycarboxylic acid is citric acid.
 - A process according to any of claims 6 to 8, in which the further metal element is silver.
- 105 10. A process according to claim 6, substantially as described in either Example.
- 11. A surgical dressing which comprises a product according to claim 4 or claim 5, or the product of a process according to claim 9 110 or claim 10.

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